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APPLICATION NO.	FILING DATE	FIRST NAMED INVENTOR	ATTORNEY DOCKET NO.	CONFIRMATION NO.
09/945,353	08/31/2001	Timothy Hla	UCT-0012-P	2675
23413	7590 03/25/2005		EXAMINER	
CANTOR COLBURN, LLP			MCGARRY, SEAN	
55 GRIFFIN ROAD SOUTH BLOOMFIELD, CT 06002			ART UNIT	PAPER NUMBER
	•		1635	

DATE MAILED: 03/25/2005

Please find below and/or attached an Office communication concerning this application or proceeding.

		Application No.	Applicant(s)				
		09/945,353	HLA ET AL.				
	Office Action Summary	Examiner	Art Unit				
		Sean R. McGarry	1635				
The MAILING DATE of this communication appears on the cover sheet with the correspondence address Period for Reply							
THE N - Exten after S - If the - If NO - Failur Any re	DRTENED STATUTORY PERIOD FOR REMAILING DATE OF THIS COMMUNICATIOnsions of time may be available under the provisions of 37 CFI SIX (6) MONTHS from the mailing date of this communication period for reply specified above is less than thirty (30) days, a period for reply is specified above, the maximum statutory perectore to reply within the set or extended period for reply will, by steply received by the Office later than three months after the mid patent term adjustment. See 37 CFR 1.704(b).	N. R 1.136(a). In no event, however, reply within the statutory minimur riod will apply and will expire SIX atute, cause the application to be	may a reply be timely filed n of thirty (30) days will be considered tim (6) MONTHS from the mailing date of this come ABANDONED (35 U.S.C. § 133).	ely. communication.			
Status							
1)⊠	1) Responsive to communication(s) filed on <u>08 August 2004</u> .						
2a)□	This action is FINAL . 2b)⊠ 1	his action is non-final.					
•	Since this application is in condition for allowance except for formal matters, prosecution as to the merits is closed in accordance with the practice under <i>Ex parte Quayle</i> , 1935 C.D. 11, 453 O.G. 213.						
Dispositi	on of Claims						
4) ☐ Claim(s) 1-12 is/are pending in the application. 4a) Of the above claim(s) 2,3,5-8 and 10-12 is/are withdrawn from consideration. 5) ☐ Claim(s) is/are allowed. 6) ☐ Claim(s) 1,4 and 9 is/are rejected. 7) ☐ Claim(s) is/are objected to. 8) ☐ Claim(s) are subject to restriction and/or election requirement.							
Application	on Papers						
10) 🖾 -	The specification is objected to by the Examember The drawing(s) filed on 31 August 2001 is/a Applicant may not request that any objection to Replacement drawing sheet(s) including the confine oath or declaration is objected to by the	re: a) \square accepted or b) the drawing(s) be held in a rection is required if the di	abeyance. See 37 CFR 1.85(a). rawing(s) is objected to. See 37 (CFR 1.121(d).			
Priority u	nder 35 U.S.C. § 119						
12) <u></u> / a)[Acknowledgment is made of a claim for fore All b) Some * c) None of: 1. Certified copies of the priority docum 2. Certified copies of the priority docum 3. Copies of the certified copies of the papplication from the International Buree the attached detailed Office action for a	ents have been receive ents have been receive priority documents have reau (PCT Rule 17.2(a)	d. d in Application No been received in this Nationa).	al Stage			
Attachment	r(s)						
2) Notice 3) Inform	e of References Cited (PTO-892) e of Draftsperson's Patent Drawing Review (PTO-948) nation Disclosure Statement(s) (PTO-1449 or PTO/SB · No(s)/Mail Date <u>3/21/02, 9/22/03</u> .	Pap /08) 5) ☐ Not	rview Summary (PTO-413) er No(s)/Mail Date ice of Informal Patent Application (P ⁻ er:	TO-152)			

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DETAILED ACTION

Applicant's election with traverse of Group III in the reply filed on 8/07/04 is acknowledged. The traversal is on the ground(s) that the resetriction is traversed. This is not found persuasive because applicant has not provided any particular arguments that show any error in the reasons set froth in the restriction of record..

The requirement is still deemed proper and is therefore made FINAL.

Claims 2, 3, 5-8, 10-12 withdrawn from further consideration pursuant to 37 CFR 1.142(b), as being drawn to a nonelected invention, there being no allowable generic or linking claim. Applicant timely traversed the restriction (election) requirement in the reply filed on 8/07/04.

Applicant is reminded that upon the cancellation of claims to a non-elected invention, the inventorship must be amended in compliance with 37 CFR 1.48(b) if one or more of the currently named inventors is no longer an inventor of at least one claim remaining in the application. Any amendment of inventorship must be accompanied by a request under 37 CFR 1.48(b) and by the fee required under 37 CFR 1.17(i).

Claims 1, 4, and 9 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the written description requirement. The claim(s) contains subject matter which was not described in the specification in such a way as to reasonably convey to

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one skilled in the relevant art that the inventor(s), at the time the application was filed, had possession of the claimed invention. This is a written description rejection.

The instant invention is broadly drawn to a method of inhibiting angiogenesis in vivo (in a whole animal) via the administration of a composition that comprises a pharmaceutically effective amount of an antagonist of EDG-1 signal transduction. The scope of potential inhibitors is quite large. For example the scope of the invention is not limited to inhibitors of EDG-1 which class of potential compounds is so large as to include any small molecule inhibitor, antibodies, antisense oligonucleotides, peptide inhibitors, etc. The scope of potential inhibitors is so vast as to include any type of antagonist compound that may antagonize EDG-1 signal transduction. This means that any component of any EDG-1 signal transduction cascade, upstream or downstream, may be a potential target of the antagonist used in the instantly claimed methods. The scope of potential antagonist compounds is indeed vast. The invention further reads on inhibiting in any animal species, for example.

The instant specification discloses two antisense oligonucleotides targeted to human EDG-1 which inhibits EDG-1 in cell in culture and suggests methods to find other potential antagonist of EDG-1 signal transduction. The specification as filed does not provide the actual structure of any other EDG-1 signal transduction, for example. If applicant believes that the specification as filed discloses other antagonist structures applicant is invited to point to such disclosure with particularity. The prior art does not provide the description lacking in the instant specification. If applicant believes that the prior art discloses sufficient structures of antagonists known in the prior art to inhibit

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EDG-1 signal transduction in vivo or which provide for inhibition of unwanted angiogenesis applicant may bring these to the examiners attention and such disclosure will be taken into consideration. The invention further includes the further limitation (in claim 4) of also including an Akt kinase inhibitor. Again the disclosure of such inhibitors is spare in comparison to the scope of that embraced within the scope of the claims.

The instant specification fails to provide a description of a sufficient number of species of antagonists that would be representative of the genus embrace for use with the instantly claimed methods. The specification fails to provide a description of any particular structure or structures that would be shared within the genus of antagonist such that one in the art would be apprised of the structure which corresponds to the function of antagonizing EDG-1 signal transduction. The specification has not shown any compounds that have been shown to inhibit angiogenesis in an animal or which has been shown to inhibit EDG-1 signal transduction in vivo. The specification provides insufficient written description to support the genus encompassed by the claim.

<u>Vas-Cath Inc. v. Mahurkar</u>, 19 USPQ2d 1111, makes clear that "applicant must convey with reasonable clarity to those skilled in the art that, as of the filing date sought, he or she was in possession of the invention. The invention is, for purposes of the 'written description' inquiry, whatever is now claimed." (See page 1117.) The specification does not "clearly allow persons of ordinary skill in the art to recognize that [he or she] invented what is claimed." (See <u>Vas-Cath</u> at page 1116.)

The skilled artisan cannot envision the detailed chemical structure of the encompassed antagonists, regardless of the complexity or simplicity of the method of

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isolation. Adequate written description requires more than a mere statement that it is part of the invention and reference to a potential method for isolating it. See Fiers v. Revel, 25 USPQ2d 1601, 1606 (CAFC 1993) and Amgen Inc. V. Chugai Pharmaceutical Co. Ltd., 18 USPQ2d 1016. In Fiddes v. Baird, 30 USPQ2d 1481, 1483, claims directed to mammalian FGF's were found unpatentable due to lack of written description for the broad class. The specification provided only the bovine sequence. University of California v. Eli Lilly and Co., 43 USPQ2d 1398, 1404, 1405 held that:

...To fulfill the written description requirement, a patent specification must describe an invention and do so in sufficient detail that one skilled in the art can clearly conclude that "the inventor invented the claimed invention." Lockwood v. American Airlines, Inc., 107 F.3d 1565, 1572, 41 USPQ2d 1961, 1966 (1997); In re Gosteli, 872 F.2d 1008, 1012, 10 USPQ2d 1614, 1618 (Fed. Cir. 1989) (" The description must clearly allow persons of ordinary skill in the art to recognize that [the inventor] invented what is claimed."). Thus, an applicant complies with the written description requirement "by describing the invention, with all its claimed limitations, not that which makes it obvious," and by using "such descriptive means as words, structures, figures, diagrams, formulas, etc., that set forth the claimed invention." Lockwood, 107 F.3d at 1572, 41 USPQ2d at 1966.

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An adequate written description of a DNA, such as the cDNA of the recombinant plasmids and microorganisms of the '525 patent, "requires a precise definition, such as by structure, formula, chemical name, or physical properties," not a mere wish or plan for obtaining the claimed chemical invention. *Fiers v. Revel*, 984 F.2d 1164, 1171, 25 USPQ2d 1601, 1606 (Fed. Cir. 1993). Accordingly, "an adequate written description of a DNA requires more than a mere statement that it is part of the invention and reference to a potential method for isolating it; what is required is a description of the DNA itself." Id. at 1170, 25 USPQ2d at 1606.

The name cDNA is not itself a written description of that DNA; it conveys no distinguishing information concerning its identity. While the example provides a process for obtaining human insulin-encoding cDNA, there is no further information in the patent pertaining to that cDNA's relevant structural or physical characteristics; in other words, it thus does not describe human insulin cDNA. Describing a method of preparing a cDNA or even describing the protein that the cDNA encodes, as the example does, does not necessarily describe the cDNA itself. No sequence information indicating which nucleotides constitute human cDNA appears in the patent, as appears for rat cDNA in Example 5 of the patent. Accordingly, the specification does not provide a written description of the invention of claim 5.

Therefore, only SEQ ID NO: XXX but not the full breadth of the claim (or none of the sequences encompassed by the claim) meets the written description provision of 35

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USC 112, first paragraph. The species specifically disclosed are not representative of the genus because the genus is highly variant. Applicant is reminded that <u>Vas-Cath</u> makes clear that the written description provision of 35 USC 112 is severable from its enablement provision. (See page 1115.)

Applicant is also directed to <u>University of Rochester v. G.D. Searle & Co.</u>, 69USPQ2d (CA FC 2004). One would not know how to make the claimed substance other than by trial and error process." "Regardless whether a compound is claimed per se or a method is claimed that entails the use of the compound, the inventor cannot lay claim to that subject matter unless he can provide a description of the compound sufficient to distinguish infringing compounds from non-infringing compounds, or infringing methods from non-infringing methods... [t]he claimed method depends upon finding a compound that's electively inhibits PGHS-2 activity. Without such a compound, it is impossible to practice the claimed method of treatment.

Claims 1, 4, and 9 are rejected under 35 U.S.C. 112, first paragraph, as failing to comply with the enablement requirement. The claim(s) contains subject matter which was not described in the specification in such a way as to enable one skilled in the art to which it pertains, or with which it is most nearly connected, to make and/or use the invention.

The claimed invention is broadly drawn to a method of inhibiting angiogenesis in vivo and methods of treating disease via inhibiting angiogenesis via the administration

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of antagonists of EDG-1 signal transduction. As was described above, the scope of inhibitors is vast. The disclosure of specific potential inhibitors is minimal. Fort example the specification discloses two antisense oligonucleotides which were shown to inhibit EDG-1 in cells in culture. The specification fails to provide any working examples that show of would show by correlation the inhibition of angiogenesis in vivo or the treatment of disease via an EDG-1 antagonist.

It is noted that the art of biotechnology is an unpredictable art and that the exemplified antisense oligonucleotides of the instant specification are part of an unpredictable art of nucleic acid therapy. It is noted that the claims are so broad to read on any type of antagonist but the following will demonstrate that even a narrow range of what is contemplated is unpredictable and not enabled.

The specification fails to provide any specific guidance for antisense based therapy other than providing two potential oligonucleotides that might be tested where there is not specific guidance for targeting or delivery in an in vivo and/or therapeutic setting.

Jen et al [STEM CELLS Vol. 18:307-319, 2000] discuss antisense based therapy and the challenges that remain before the use of antisense becomes routine in a therapeutic setting. Jen et al discuss the advances made in the art but also indicate that progress needs to be made in the art. In the conclusion of their review Jen et al assert "[g]iven the state of the art, it is perhaps not surprising that effective and efficient clinical translation of the antisense strategy has remained elusive." It is also stated "[t]he key challenges to this field have been outlined above. [l]t is clear that they will have to be solved if this approach to specific antitumor therapy is to become a useful treatment

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approach. [a] large number of diverse and talented groups are working on this problem, and we can all hope that their efforts will help lead to establishment of this promising form of therapy." It is clear from Jen et al that the state of the art of antisense is unpredictable and those highly skilled in the art are working towards making the art of antisense therapy more predictable but have many obstacles to overcome.

Branch [TIBS Vol. 23, February 1998 and cited by applicant on form 1449 filed 3/21/02] addresses the unpredictability and the problems faced in the antisense art with the following statements: "[a]ntisense molecules and ribozymes capture the imagination with their promise or rational drug design and exquisite specificity. [h]owever, they are far more difficult to produce than was originally anticipated, and their ability to eliminate the function of a single gene has never been proven."; "[t]o minimize unwanted nonantisense effects, investigators are searching for antisense compounds and ribozymes whose targets sites are particularly vulnerable to attack. [t]his is a challenging guest.": "[h]owever, their unpredictability confounds research applications of nucleic acid reagents."; "[n]on-antisense effects are not the only impediments to rational antisense drug design. [t]he internal structures of target RNAs and their associations with cellular proteins create physical barriers, which render most potential binding sites inaccessible to antisense molecules."; "Years of investigation can be required to figure out what an 'antisense' molecule is actually doing...."; "Because knowledge of their underlying mechanism is typically acting, non-antisense effects muddy the waters."; "because biologically active compounds generally have a variety of effects, dose-response curves are always needed to establish a compounds primary pharmacological identity.

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[a]ntisense compounds are no exception. [a]s is true of all pharmaceuticals, the value of a potential antisense drug can only be judged after its intended clinical use is known, and quantitative information about its dose-response curve and therapeutic index is known."; [c]ompared to the dose response curves of conventional drugs, which typically span two to three orders of magnitude, those of antisense drugs, extend only across a narrow concentration range."; "[b]ecause it is very difficult to predict what portions of an RNA molecule will be accessible *in vivo*, effective antisense molecules must be determined empirically by screening large number of candidates for their ability to act inside cells."; "[b]inding is the rare exception rather than the rule, and antisense molecules are excluded from most complementary sites. [s]ince accessibility cannot be predicted, rational design of antisense molecules is not possible."; and, "[t]he relationship between accessibility to ODN binding and vulnerability to ODN-mediated antisense inhibition *in vivo* is beginning to be explored. . . [i]t is not yet clear whether *in vitro* screening techniques. . . will identify ODNs that are effective *in vivo*."

One in the art would be required to perform an undue quantity of trial and error experimentation to determine potential antagonists from the vast range contemplated and then to further determine how to use such compounds in a method of treatment where it has been demonstrated that at least for antisense oligonucleotides one in the art requires specific guidance on methods or targeting and delivery, for example.

The type of experimentation required to practice the invention more broadly than it is exemplified is a factor in the enablement analysis, but is not dispositive. In this

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case, the more or less standard nature of each type of experiment to find potential antagonists required to expand the scope of the enabled invention is outweighed by the sheer quantity of experimentation required to practice the full scope of the claims, the unpredictability of the art generally and the claimed method in particular, and the lack of guidance in the specification regarding the direction in which the experimentation should proceed.

The prior art made of record and not relied upon is considered pertinent to applicant's disclosure. The following references are not relied upon in the Official Action but are considered to be pertinent to applicants' disclosure. US 6,323,333; US 6,423,508; US 6,649,640; and US 6,777,439.

Any inquiry concerning this communication or earlier communications from the examiner should be directed to Sean R. McGarry whose telephone number is (571) 272-0761. The examiner can normally be reached on M-Th (6:00-4:30).

If attempts to reach the examiner by telephone are unsuccessful, the examiner's supervisor, John LeGuyader can be reached on (571) 272-0760. The fax phone number for the organization where this application or proceeding is assigned is 703-872-9306.

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Sean R McGarry Primary Examiner Art Unit 1635

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